PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISH	HED 1	DER THE PATENT COOPERATION	TREATY (PCT)
(51) International Patent Classification 7:		1) International Publication Number:	WO 00/48957
C03C 25/26, C08J 5/08	A1	3) International Publication Date: 24	August 2000 (24.08.00)
 (21) International Application Number: PCT/US (22) International Filing Date: 11 February 2000 ((30) Priority Data: 09/250,720 16 February 1999 (16.02.99) (71) Applicant (for all designated States except US): CORNING [US/US]; One Owens Corning Parkway OH 43659 (US). (72) Inventors; and (75) Inventors/Applicants (for US only): CAMPBELL, [US/US]; 3020 Brackeberry Drive, Anderson, S (US). VICKERY, Eric, L. [US/US]; 255 Pauli Williamston, SC 29697 (US). (74) Agents: ECKERT, Inger, H. et al.; Owens Corning S Technology Center, 2790 Columbus Road, Buildi Granville, OH 43023-1200 (US). 	OWEN, Toled Les, C 296 an Roa	(81) Designated States: AE, AL, AM, AT, BR, BY, CA, CH, CN, CR, CU, C ES, FI, GB, GD, GE, GH, GM, HR, KE, KG, KP, KR, KZ, LC, LK, LR, MD, MG, MK, MN, MW, MX, NO, SD, SE, SG, SI, SK, SL, TI, TM, US, UZ, VN, YU, ZA, ZW, ARIPO LS, MW, SD, SL, SZ, TZ, UG, ZW, AZ, BY, KG, KZ, MD, RU, TI, TM, BE, CH, CY, DE, DK, ES, FI, FR, MC, NL, PT, SE), OAPI patent (BF GA, GN, GW, ML, MR, NE, SN, T Published With international search report. Before the expiration of the time is claims and to be republished in the amendments.	Z, DE, DK, DM, EE, HU, ID, IL, IN, IS, IP, LS, LT, LU, LV, MA, NZ, PL, PT, RO, RU, TR, TT, TZ, UA, UG, patent (GH, GM, KE, Le Corona patent (AT, GB, GR, IE, IT, LU, BJ, CF, CG, CI, CM, D, TG).
(54) Title: SIZING COMPOSITION FOR GLASS FIB MATRIX POLYMERS	ERS U	D TO REINFORCE THERMOPLASTIC O	R THERMOSETTING
(57) Abstract			
The present invention relates to a sizing composition particularly, polypropylene. The composites produced with	for gla	fibers used to reinforce thermoplastic resins, s	uch as polyolefins, and

The present invention relates to a sizing composition for glass fibers used to reinforce thermoplastic resins, such as polyolefins, and particularly, polypropylene. The composites produced with the fiber strands coated with such sizing compositions have improved fiber adhesion and retention of fiber length. In addition, a method for improving tensile creep and tensile fatigue in polypropylene composites reinforced with glass fibers is provided.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL .	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT .	Austria	FR	France	LU	Luxembourg	SN	Senegal
	Australia	GA	Gabon	LV	Larvia	SZ	Swaziland
AZ .	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
	Barbados	GH	Ghana	MG	Madagascar	ТJ	Tajikistan
	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
BJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
	Congo	KE	Кепуа	NL	Netherlands	YU	Yugoslavia
CH :	Switzerland	KG	Kyrgyzstan	NO	Norway	zw	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE .	Germany	u	Liechtenstein	SD	Sudan		
	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		

SIZING COMPOSITION FOR GLASS FIBERS USED TO REINFORCE THERMOPLASTIC OR THERMOSETTING MATRIX POLYMERS

TECHNICAL FIELD AND INDUSTRIAL

5 APPLICABILITY OF THE INVENTION

The present invention relates to a sizing composition for glass fibers used to reinforce thermoplastic resins, such as polyolefins, and particularly, polypropylene. The composites produced with the fiber strands coated with such sizing compositions have improved fiber-matrix adhesion and retention of fiber length, and exhibit improved tensile creep and tensile fatigue.

BACKGROUND OF THE INVENTION

10

15

20

25

30

The composite industry has been using glass fibers in various forms for reinforcing polymeric matrices, concrete, paper, paper/foil and cables to produce a variety of products. Glass fibers have been used in the form of continuous or chopped filaments, strands and rovings, as well as woven, non-woven and braided fabrics, and meshes and scrims as reinforcements.

Production of glass fibers for the composite market typically involves attenuation of the glass fibers from molten streams of fiberizable glass material emanating from a bushing or like device connected to a furnace containing molten fiberizable glass material. The glass fibers are attenuated by conventional means such as winders and choppers. In the process of producing glass fibers, a chemical composition is usually applied to them shortly after they are attenuated from molten streams of glass. This chemical composition, or sizing, is typically an aqueous mixture containing one or more starches or non-starch film formers, one or more lubricants and emulsifiers, and optionally, one or more organo functional silanes. This chemical composition or sizing is necessary in order to retard abrasion of the glass fibers when they are gathered into glass fiber strands, as well as in subsequent processing, such as winding, twisting, weaving and braiding of the glass fibers into the final product. In addition to acting as a processing aid to prevent fiber damage from handling, the size composition is typically designed to provide the composite with improved physical properties, hydrolytic stability and corrosion resistance. After application of the sizing, the fibers are then dried either in the package form or in the chopped strand form before they are used as reinforcement.

Although numerous glass fiber sizing compositions are known, a continuing need exists for new sizing compositions that provide improved matrix resin adhesion and retention of fiber length when chopped glass fibers coated with such sizing composition are used in the reinforcement of polymeric matrices, particularly, polypropylene. This need is fulfilled by the sizing composition and process of the invention described below.

SUMMARY OF THE INVENTION

5

10

15

20

25

30

According to this invention, an improved sizing composition is provided for treating glass fibers used to reinforce thermoplastic or thermosetting matrix polymers. The sizing compositions of the present invention are particularly well suited for use in connection with the reinforcement of polyolefin matrices, e.g., polypropylene, and provide such reinforced composites with improved tensile creep and tensile fatigue. In particular, the sizing composition provides better adhesion between the glass fiber surface and matrix resin, improved nucleation in the polypropylene at the interface region between the sizing and the matrix, and increased retention of filament length in the resulting molded parts.

According to one aspect of the invention, the sizing composition comprises: (a) a cationic high molecular weight acid or anhydride modified polypropylene film former, (b) an amino silane coupling agent, (c) a cationic lubricant, and (d) a nucleating agent. Preferably, the high molecular weight polypropylene film former has been modified by grafting maleic anhydride thereto, and has a molecular weight in excess of 35,000 Daltons, and the nucleating agent is preferably a C₁₀-C₁₈ fatty acid. The combination of these materials in the sizing composition provides improved adhesion and retention of fiber length when chopped glass fibers coated with such sizing composition are used in the reinforcement of polypropylene matrices.

Moreover, another aspect of this invention is directed to a process for improving the tensile creep and tensile fatigue exhibited by such composites under loads by increasing the adhesion between the glass fiber surface and matrix resin, nucleation in the polypropylene at the interface region between the sizing and the matrix, and retained filament length in the molded parts. The process involves reinforcing a polymeric matrix using glass fibers treated with the sizing compositions disclosed above.

DETAILED DESCRIPTION AND PREFERRED EMBODIMENTS OF THE INVENTION

5

10

15

20

25

30

The glass fiber sizing compositions of the present invention comprise: (a) a cationic high molecular weight acid or anhydride modified polypropylene film former, (b) an amino silane coupling agent, (c) a cationic lubricant, and (d) a nucleating agent.

As used herein, the terms "size" or "sizing" refer to a composition applied to glass fibers shortly after the glass fibers are formed. The glass fibers sized in accordance with the present invention are compatible with thermoplastic matrix polymers such as polyolefin matrices and in particular, polypropylene. Polypropylene reinforced with the sized glass fibers of the present invention exhibits improved fiber adhesion and retention of fiber length, which provides composites with reduced tensile creep and fatigue under loads.

Numerous film forming polymers, including carboxylic acid or anhydride modified polyolefins, have been conventionally used in sizes for glass fibers. See, for example, U.S. Patent Nos. 5,648,169 and 5,646,207. However, the film formers used in the present invention are cationic high molecular weight acid or anhydride modified polypropylenes that have a molecular weight greater than 35,000, and preferably, greater than 80,000 Daltons. These include emulsions of polypropylenes, mixtures of polypropylenes, or polypropylene derivatives, chemically modified to graft thereto maleic anhydride. Suitable polypropylene film formers for use in the present sizing composition include PP448 and PP353C commercially available from E.I. DuPont de Nemours; PB3001, PB3150, PB3002 commercially available from Uniroyal (Polybond Group); and G-3015 commercially available from Eastman Chemical. These polypropylenes have molecular weights ranging from approximately 35,000 to over 80,000, and maleation (maleic acid/anhydride grafting)ranging from less than 0.5 weight percent to approximately 1.8 weight percent. In a preferred embodiment, DuPont PP353C maleated polypropylene is used as the film former in the sizing compositions of the present invention. The film former of the present invention is preferably present in an amount of from about 40 percent to about 80 percent of the total dried sizing.

In addition, other conventional film formers may be present in the sizing composition in addition to the polypropylene film former. For example, polyurethane film formers may be useful in the sizing composition of the present invention. One such

polyurethane film former is Aquathane 52-00-01 commercially available from Nippon Ink & Chemicals of Japan.

The aminosilane coupling agents that may be used in accordance with the present invention include any of those that are conventionally used for such purposes. For example, a mono- or di-aminated aminosilane such as gamma-aminopropyltriethoxysilane or a N-beta-(aminoethyl)-gamma-aminopropyltrimethoxysilane or any other similar aminosilane that is commercially available may conveniently be used in the present invention. For example, U.S. Patents Nos. 5,470,658 and 5,646,207 describe suitable aminosilanes useful in sizing compositions. Preferably, gamma-

aminopropyltriethoxysilane, which is commercially available from Osi Specialties, Tarrytown, N.Y. under the trade designation A-1100, is used in the present invention. The aminosilane is preferably present in an amount of from about 10 to about 20 percent of the total dried sizing.

10

15

20

25

30

The cationic lubricants used in the sizing compositions of the present invention reduce filament damage during the fiber forming and chopping processes. Numerous cationic lubricants are known in the art for use in sizing compositions to treat glass fibers. For example, stearic ethanolamide (Lubsize K12 commercially available from Alpha Owens Corning), or polyethyleneimine polyamides (commercially available from Henkel Inc. under the trade designations Emery 6760 and Emery 6717) can be used in the sizing compositions of the present invention. The cationic lubricant is preferably present in the sizing composition in an amount of from about 0.25 to about 1.3 percent of the total dried sizing.

The nucleating agents used in accordance with the present invention are fatty acids which provide two beneficial attributes to the sized glass fibers. First, the fatty acid provides lubricity to the glass fiber in order to minimize filament length degradation during the high shear forces of compounding and molding. Second, the fatty acid acts as a nucleating agent to affect crystalline growth of the polypropylene matrix, specifically at the interface region. In particular, the nucleating agent of the present invention is preferably a C₁₀-C₁₈ fatty acid. An example of a suitable nucleating agent is a fatty acid blend such as Moldpro 932 commercially available from Witco. The nucleating agent is preferably present in the sizing composition in an amount of from about 8.0 to about 15.0 percent of the total dried sizing.

The pH of the sizing composition is generally basic and is preferably greater than

about 8.5. It may be necessary to add a pH buffer in order to maintain the pH since the cationic components present in the sizing composition may destabilize the sizing composition pH.

In addition to the above components, other components that are normally added to glass fiber sizing composition can also be present. For example, biocides, anti-foaming agents, plasticizers, humectants, antistatic agents, flame retardants, and uv light stabilizers may be used in the sizing compositions.

5

10

15

20

25

30

The glass fibers employed in the present invention can be any of the fibers known in the art for such purpose. For example, suitable fibers are typically fibers having a diameter of from about 9 to about 20 microns, and suitable strands generally contain from about 2000 to about 5000 fibers. Preferably, the fibers are in the range of from about 10 to about 17 microns and the strands formed contain from about 2400 to about 4100 fibers. The glass fibers used in the context of this invention are generally intended to reinforce thermoplastic matrices such as polyolefin matrices and, preferably, polypropylene matrices.

Shortly after being formed, the glass fibers may be treated with the sizing composition of the present invention according to methods well known in the art. The size can generally be applied with sprayers, rollers, belts, metering devices, hot melt applicators, or other similar application devices. The sized fibers are then gathered into bundles or strands comprising a plurality of individual fibers. After their formation and treatment, the strands are generally wound onto a spool or forming package and also may be chopped. The forming packages or chopped strands are usually dried either in an oven or at room temperature to remove some of the moisture from the fibers.

After application of the sizing, the size present on the fibers may be measured by determining the loss on ignition (LOI) of the rovings. The LOI refers to the reduction in weight of the fibers after heating by burning or pyrolyzing to remove the organic size from the fibers. Preferably, the sized glass fibers of the invention have an LOI of from about 0.5 percent to about 1.1 percent.

The dried fibers treated with the composition described above may be used to reinforce thermoplastic matrix polymers such as polyolefins and, particularly, polypropylene, in any of the molding processes known to those skilled in the art utilizing either chopped strand, continuous strand, or a mixture thereof. Preferably, the molding process uses chopped fiber strands mixed or compounded with matrix polymer pellets,

e.g., polypropylene, to produce a homogeneous batch of molding compound. The batch can then be used, for example, in compression or injection molding to produce the fiber reinforced polymeric part or shaped device.

In another embodiment of the invention, this invention relates to a process of improving the tensile creep and tensile fatigue of such molded parts under loads. This is accomplished by increasing the adhesion between the glass fiber surface and matrix resin, nucleation in the polypropylene at the interface region between the sizing and the matrix, and retained filament length in the molded parts. The process involves reinforcing a polymeric matrix with glass fibers treated with the sizing compositions disclosed above.

5

10

In order that the invention may be more readily understood, reference is made to the following examples, which are intended to be illustrative of the invention, but are not intended to be limiting in scope.

EXAMPLE I

This example illustrates the preparation of preferred sizing compositions for glass fibers according to the present invention. In Table 1, there are shown various sizing compositions of the invention, samples 1-6, for treating glass fibers.

TABLE 1

SAMPLE	1			2		3
INGREDIENTS	%	Amount in grams	%	Amount in grams	%	Amount in grams
Al 100 ¹ (silane)	1.5	150	1.5	150	1.5	150
Moldpro 932 ²	3.0	300	3.0	300	3.0	300
PP448C ³	11.0	1100	11.0	1100	11.0	1100
Aquathane 52-00-01	0.5	50	0.5	50	0.5	50
K-12 ⁵ (8.8%)	0.08	8	0.24	24	-	-
6760T ⁶ (50%)		•	-	-	0.01	1
Strand solids		0.64		0.57		0.58
WATER	83.92	8,021	83.76	7,957	83.99	8,819
Quantity		9,558		9,500		10,500

A1100 is a gamma-aminopropyltriethoxysilane obtained from Osi Specialties, Tarrytown, N.Y.

Moldpro 932 is a fatty acid nucleating agent obtained from Whitco.

³ PP448C is a high molecular weight maleonated polypropylene obtained from E.I. DuPont de Nemours.

⁴ Aquathane 52-00-01 is a polyurethane film former obtained from Nippon Ink & Chemicals of Japan.

Lubsize K-12 is a cationic lubricant commercially available from Alpha Owens Corning.
 6760 T is a polyethylene polyamide salt cationic lubricant obtained from Henkel, Inc.

SAMPLE		4		5		6	
INGREDIENTS	%	Amount in grams	%	Amount in grams	%	Amount in grams	
Al 100 ¹ (silane)	1.5	150	1.5	150	1.5	150	
Moldpro 9321	3.0	300	3.0	300	3.0	300	
PP448C ¹	12.0	1200	12.0	1200	12.0	1200	
Aquathane 52-00-01	0.5	50	0.5	50	0.5	50	
K-12 ¹ (8.8%)	0.08	8	0.24	24	-	-	
6760T ¹ (50%)	-	-	-	-	0.01	1	
Strand solids		0.97		1.07			
WATER	82.92	7,877	82.76	8,251	82.99	7,884	
Quantity		9,500		9,970		9,500	

The sizing composition of samples 1-6 were formulated in the following manner. A premix of the polypropylene dispersed in water was first made. To this premix, is added the fatty acid emulsion and the polyurethane dispersion under agitation. A separate premix of the organosilane in water was prepared and added to the former premix. Finally, additional deionized water was added to attain the desired solids content.

Glass fibers having a diameter of 13.7 microns were treated with the sizing compositions 1-6 and chopped into lengths suitable for use as a reinforcement media in molded parts. Test specimens were prepared by compounding the sized fiber segments with a polypropylene resin, comprising 5% by weight PB3002 maleated polypropylene and 95% by weight Montel 6523 homopolypropylene resin, in a 2-1/2 inch single screw HPM extruder, and injection molding the resulting compound into plaques in a 150 ton Cincinnati Molding Machine. The physical properties of the resulting plaques were determined according to the procedures set forth in ASTM D638 (tensile strength); ASTM D256-81 (flexural strength); and ASTM D790 (IZOD impact strength). The

10

ASTM D256-81 (flexural strength); and ASTM D790 (IZOD impact strength). The mechanical properties of the composites reinforced with these treated fibers are set forth in Table 2.

TABLE 2

SAMPLE	TENSILE	FLEXURAL	UNNOTCHED IZOD	GLASS
	STRENGTH	STRENGTH	IMPACT STRENGTH	CONTENT
	(psi)		(ft-lb/in)	WEIGHT %
1	12,102		12.193	30.4
2	12,453		13.57	30.5
3	12.696	excellent	13.14	30.2
4	12,678	excellent	13.544	29.9
5	12,701	excellent	13.325	29.7
6	12,773	excellent	13.605	29.6

Applicants observed very good tensile and impact strengths. In particular, the addition of K-12 and 6760T (both cationic lubricants) in combination with the fatty acid nucleating agents and high molecular weight maleonated polypropylene was very beneficial in minimizing filament damage during the chopping process. It was expected that good creep and fatigue properties could be correlated with this data as determined by measuring retained filament length in the resulting molded parts.

10 <u>EXAMPLE II</u>

5

15

In order to determine how to stabilize the Moldpro 932 emulsion upon the addition of A1100 to the polypropylene emulsion mixture, the following experiments were performed. Deionized water having a pH of 7.3 was added to A1100 premix having a pH of 10.7. The pH of the A1100 solution was then adjusted according to Table 3 prior to adding the polypropylene emulsion. Then about 3 percent by weight Moldpro 932 emulsion having a pH of 4.5-5.0 was added to the main mix tank. The resulting level of precipitation of the Moldpro 932 is shown below in Table 3.

TABLE 3

A1100 PH PRIOR TO PP EMULSION ADDITION	RESULTS
10.7	Moldpro 932 dispersion; heavy precipitation
6.7 (acetic acid adjusted)	Moldpro 932 dispersion; precipitates, but not as much
5.5 (acetic acid adjusted)	slight precipitation

EXAMPLE III

Several other sizing compositions of the present invention were prepared. These compositions were made in a similar manner as those made in Example I. However, a different type of polypropylene emulsion, PP353C from DuPont, was used in some of the formulations. Table 4 describes the sizing compositions prepared using the PP 448C polypropylene emulsion (samples 7-15)and Table 5 describes the sizing compositions prepared using the PP353C polypropylene emulsion (samples 16-24).

TABLE 4

SAMPLE	7			8		9	
INGREDIENTS	%	Amount in grams	%	Amount in grams	%	Amount in grams	
A1100	1.5	225	1.5	225	1.5	225	
Acetic Acid		pH=5.1		pH=5.1		pH=5.1	
PP448C emulsion	10.0	1500	12.0	1800	12.0	1800	
Aquathane 52-00-01	0.5	75	0.0		0.5	75	
Moldpro 932 (26%)	5.0	750	5.0	750	5.0	750	
Theor. Mix Solids		4.9%		5.2%		5.4%	
Actual Quantity					-	31170	
Solids(gms) Percent Solids		14,000 4.96%		14,000 4.96%		14,000 5.33%	

10			11	12	
%	Amount in grams	%	Amount in grams	%	Amount in grams
1.5	225	1.5	225	1.5	225
	pH=5.1		pH=5.1		PH=5.1
12.0	1800	12.0		10.0	1500
0.5	75				
1.0	150	1.0	150	1.0	150
	4.36%		4.16%		3.66%
	14,000		14,000		14,000
	1.5 12.0 0.5	% Amount in grams 1.5 225 pH=5.1 12.0 1800 0.5 75 1.0 150 4.36%	% Amount in grams % 1.5 225 1.5 pH=5.1 12.0 1800 12.0 0.5 75 1.0 150 1.0 4.36% 1.0	% Amount in grams % Amount in grams 1.5 225 1.5 225 pH=5.1 pH=5.1 pH=5.1 12.0 1800 12.0 1800 0.5 75 1.0 150 1.0 150 4.36% 4.16% 14,000 14,000	% Amount in grams % Amount in grams % 1.5 225 1.5 225 1.5 pH=5.1 pH=5.1 pH=5.1 12.0 1800 10.0 0.5 75 1.0 150 1.0 150 1.0 4.36% 4.16% 4.16%

10

5

SAMPLE	13			14	15		
INGREDIENTS	%	Amount in grams	%	Amount in grams	%	Amount in grams	
A1100	1.5	225	1.5	225	1.5	225	
Acetic Acid		pH=5.1		pH=5.1		pH=5.1	
PP448C emulsion	10.0	1500	10.0	1500	11.0	1650	
Aquathane 52-00-01	0.5	75			0.25	37.5	
Moldpro 932 (26%)	1.0	150	5.0	750	3.00	450	
Theor. Mix Solids		3.86%		4.7%		4.53%	
Actual Quantity Solids (gms) Percent Solids		14,000 4.06%		14,000 4.69%		14,000 4.69%	

TABLE 5

SAMPLE	16			17	1	18	
INGREDIENTS	%	Grams	%	Grams	%	Grams	
A1100	1.5	112	1.5	112	1.5	112	
Acetic Acid		50		50		50	
PP353C emulsion	10.0	750	12.0	900	12.0	900	
Aquathane 52-00-01	0.5	37.5	-	•	0.5	37.5	
Moldpro 932 (26%)	5.0	375	5.0	375	5.0	375	
Strand Solids				0.98	1	1.05	
Actual Quantity (gms)		5,000		5,000	†	5,000	

SAMPLE	19			20	21	
INGREDIENTS	%	Grams	%	Grams	%	Grams
A1100	1.5	112	1.5	112	1.5	112
Acetic Acid		50		50		50
PP353C emulsion	12.0	900	12.0	900	10.0	750
Aquathane 52-00-01	0.5	75	-	•	-	
Moldpro 932 (26%)	1.0	75	1.0	75	1.0	75
Strand Solids		0.70		0.69		0.68
Actual Quantity (gms)		5,000		5,000		5,000

SAMPLE 22 **INGREDIENTS** % Grams Grams % Grams A1100 1.5 112 1.5 112 1.5 112 Acetic Acid 50 50 50 PP353C emulsion 10.0 750 10.0 750 11.0 825 Aquathane 52-00-01 0.5 37.5 0.25 19 Moldpro 932 (26%) 1.0 75 5.0 310 3.00 225 Strand Solids 0.7 0.74 0.87 Actual Quantity (gms) 5,000 5,000 5,000

5

10

15

EXAMPLE IV

The sizing compositions comprising the emulsion of polypropylene PP353C as shown in Table 5 were applied to glass fibers having a diameter of 9 microns according to known methods. The physical strand properties of the sized glass fibers were assessed and the results obtained therefrom are described in Table 6. The applicator speed is recorded in feet per minute (fpm), and the strand moisture content and strand solids are recorded in percent by weight of the resulting sized strands. The amount of fines present after chopping the strands into segments was determined by mechanically sieving a given amount of chopped strands through a specified opening screen for a predetermined amount of time. The accumulated fine fibers that have passed through the sieve are then weighed and recorded as a percentage by weight of the chopped strands.

The strand integrity of the dry chopped strands provides an indication of how well the glass reinforcement will resist fuzzing during glass feeding or blending. A given amount of chopped strands is subjected to agitation for a predetermined time in the presence of steel balls using a paint shaker. The resulting material is then sieved to remove fines, weighed, and recorded as a percentage of the chopped strands.

5

10

15

The long fibers in the chopped strands, which are at least twice the length of the nominal chopped length, is determined by mechanically sieving a predetermined amount of chopped strands through a vibrating screen having specified openings. The strands that do not pass through the screen are collected, weighed, and recorded as a percentage of the chopped strand segments.

TABLE 6

APPLICATOR SPEED	SAMPLE	MOISTURE (%)	SOLIDS (%)	FINES (%)	3 MIN S.I. (%)	LONG FIBER
54 fpm	21	0	0.68	2.7	3.37	(%) 0.647
52 fpm	17	0.017	0.98	2.82	17.07	2.914
52 fpm	16		PL	2.714		
52 fpm	23	0.013	0.74	3.4	46.63	1.35
52 fpm	22	0	0.7	2.32	15.24	0.797
50 fpm	24	0.093	0.87	4.5	50.9	1.42
48 fpm	18	0.021	1.05	2.85	17.8	3.35
48 fpm	20	0	0.69	2.95	21.76	0.41
48 fpm	19	0.003	0.70	2.5	12.1	0.24

Glass fiber reinforced composites were prepared as described above in Example I using the glass fibers sized with the sizing compositions detailed above. The composites were evaluated for tensile strength according to ASTM D638 and impact strength according to ASTM D790, and the results obtained are provided in Table 7.

TABLE 7

SAMPLE	TENSILE	UNNOTCHED	GLASS
	(PSI)	IZOD	CONTENT
		(FTLB/IN)	(%)
16	12,587	12.851	30
21	11,091	11.36	30.2
17	11,864	12.719	30
23	11,177	11.842	30.1
22	10,983	11.316	30.1
24	10,857	11.272	30.1
18	11,552	12.149	30.1
20	11,058	11.754	29.9
19	10,862	11.711	29.9

EXAMPLE V

Another set of experiments were conducted to confirm the effect of the cationic lubricant addition to the polypropylene sizing chemistry. This comparative experiment examined the effects of two different polypropylene emulsions with and without the cationic lubricant. Sizing compositions (samples 25-31) were prepared in accordance with the formulations provided in Table 8 and by the methods similar to those used to prepare the compositions in Example I.

TABLE 8

SAMPLE	25			26		27		28	
INGREDIENT	%	gms	%	Gms	%	Gms	%	gms	
A1100	1.5	150	1.50	150	1.5	150	1.5	150	
TA22 ⁷	225	225	2.25	225	2.25	225	2.25	225	
NH₄oH	1.4	140	1.4	140	1.4	140	1.4	140	
PP448C	11.0	1,100	11.0	1,100					
PP353C					11.0	1,100	11.0	1,100	
Aquathane 52-00-01	0.5	50	0.5	50	0.5	50	0.5	50	
K-12 (8.8%)	0.24	24			0.24	24			
6760 (50%)			0.05	5			0.05	5	
Quantity Solids		10,000		10,000		10,000		10,000	

10

15

5

SAMPLE	29			30	31	
INGREDIENT	%	gms	%	gms	%	gms
A1100	1.5	150	1.50	150	1.5	150
TA22	225	225	2.25	225	2.25	225
NH₄oH	1.4	140	1.4	140	1.4	140
PP448C	11.0	1,100				
PP353C			11.0	1,100	11.0	1,100
Aquathane 52-00-01	0.5	50	0.5	50	0.5	50
K-12 (8.8%)	0.24	24			0.24	24
6760 (50%)					0.05	5
Acetic Acid					1.0	100
Quantity Solids		10,000		10,000		10,000

Glass fibers having a diameter of 13.7 and 17 microns, respectively, were coated with the sizing compositions of samples 25-31 according to known methods, and the physical strand properties of the sized glass fibers were determined as described above in Example IV. The average values are set forth in Table 9.

⁷ TA22 is terephthalic acid commercially available from Amoco.

TABLE 9

SAMPLE I.D.	MOIST	SOLIDS	3 MIN S.I.
25	0.023	0.77	0.98
26	0.029	0.75	0.84
27	0.030	0.77	0.06
28	0.026	0.81	0.01
29	0.023	0.65	1.12
30	0.020	0.64	0.54

Based on these results, the inclusion of the cationic lubricant appears to increase
the amount of the sizing composition picked up by the fibers from the applicator.

Glass fiber reinforced composites formed from chopped glass fibers sized with the sizing compositions detailed above and a polypropylene resin according to the method described above in Example I were assessed for tensile and impact strengths. The composite properties were determined and are provided in Table 10.

10

15

TABLE 10

SAMPLE	TENSILE ST (PSI)	UNNOTCHED IZOD (FT-LB/IN)	GLASS CONTENT
25	12,175	10.140	(%)
		12.149	29.8
26	12,183	12.965	29.8
27	12,325	13.193	30
28	12,297	12.412	30
29	12,196	12.956	30
30	12,276	12.544	29.8

These results demonstrate the effect of the sizing compositions of this invention on the resulting composites. As can be readily ascertained, the sizing compositions of the invention provide excellent results when used in the reinforcement of polypropylene matrices.

EXAMPLE VI

Another set of sizing compositions were prepared and used to size glass fibers having a diameter of 13.7 microns according to known methods. The sizing compositions and the resulting solids content are provided in Table 11.

TABLE 11

SAMPLE	(32		33		34	1	35
INGREDIENTS	%	gms	%	gms	%	gms	%	gms
A1100	1.83	183	1.16	116	1.5	150	1.16	116
PP353C	14.66	1,466	9.36	936	12.0	1200	9.36	936
Aquathane 52-00-01	0.61	61	0.39	39	0.5	50	0.39	39
K-12 (8.8%)	0.29	29	0.19	19	0.24	24	0.24	24
Moldpro 932	3.67	367	2.34	234	3.0	300	2.34	234
Acetic Acid				pH 7.0		pH 8.5		
Quantity Solid (gms) Percent Solids (%) PH		10,000 5.73 6.8		10,000 3.67 8.7		10,000 4.72 6.7		10,000 3.68 9.5

SAMPLE		36		37		38		39
INGREDIENTS	%	gms	%	gms	%	gms	%	gms
A1100	1.16	116	1.83	183	1.83	183	1.16	116
PP353C	9.36	936	14.66	1,466	14.66	1,466	9.36	936
Aquathane 52-00-01	0.39	39	0.61	61	0.61	61	0.39	39
K-12 (8.8%)					0.29	29	0.24	24
Moldpro 932	2.34	234	3.67	367	3.67	367	2.34	234
Acetic Acid				pH 7.0		pH 7.0		pH 7.0
Quantity Solids (gms) Percent Solids (%) pH		10,000 3.71 7.6		10,000 5.76 6.4		10,000 5.84 6.4		10,000 3.76 6.5

SAMPLE	40			41	42	
INGREDIENTS	%	gms	%	gms	%	gms
A1100	1.83	183	1.5	150	1.5	150
PP353C	14.66	1,466			12.0	1,200
Aquathane 52-00-01	0.61	61	0.5	50	0.5	50
K-12 (8.8%)	0.29	29	0.24	24	0.24	24
Moldpro 932	3.67	367	3.0	300	3.0	300
Acetic Acid						
			12.0	1,200		
Quantity Solids (gms) Percent Solids (%) pH		10,000 5.86 6.6		10,000 4.86 10.3		10,000

5

In the method of preparing these compositions, it was noted that when the PP353C polypropylene emulsion was added to the A1100 premix, the mixture tended to precipitate. Therefore, the pH was adjusted to near neutral with acetic acid in order to stabilize the mixture. The addition of the PP448C polypropylene emulsion to the A1100

premix, on the other hand, was stable.

In addition, it was noted that the stability of the sizing with compositions containing the 353C polypropylene emulsion was even more stable when the A1100 premix was diluted with water at a ratio of 10 parts of water to 1 part A1100.

EXAMPLE VII

The sizing compositions of Samples 32-41 were used to size glass fibers having a diameter of 13.7 microns according to known methods. The moisture, solids content, and strand integrity of the sized fiber strands were determined as detailed above in Example IV, and are set forth in Table 12.

10

5

TABLE 12

SAMPLE	MOIST (%)	SOLIDS (%)	3 MIN. S.I. (%)
32	0.032	0.67	0.15
33	0.022	0.46	1.89
34	0.027	0.49	0.06
35	0.016	0.40	4.36
36	0.009	0.40	4.05
37	0.018	0.71	0.83
38	0.023	0.74	0.58
39	0.012	0.44	6.56
40	0.009	0.71	0.61
41	0.015	0.49	0.49

The sized fibers were then used to prepare composites according to the methods described above in Example I. The properties of the resulting composites prepared from these sized fibers are provided in Table 13.

15

TABLE 13

SAMPLE	TENSILE ST	UNNOTCHED IZOD	GLASS	STRAND
	(PSI)	(FT-LB/IN)	CONTENT	SOLID
			(%)	(%)
32	12,800	13.333	30.2	0.67
33	12,744	13.026	30.1	0.46
34	12,720	13.772	30.3	0.49
35	12,448	13.553	30.1	0.4
36	12,316	13.465	29.9	0.4
37	12,273	13.772	30.1	0.71
38	12,479	13.772	29.8	0.74
39	12,518	13.289	30.3	0.44
40	12,422	13.772	29.9	0.71
41	12.053	13.947	30.6	0.49

Furthermore, the composites formed with the fibers sized with the compositions of samples 32-41 were tested for tensile fatigue according to the procedures of ASTM D-2991. The results of this analysis are set forth in Table 14, in terms of the average number of cycles until failure occurred at 8.4 Kpsi.

TABLE 14

SAMPLE	TENSILE FATIGUE AT 8.4 KPSI
32	1,384
33	1,500
34	1,428
35	1,438
36	1,589
37	1,998
38	1,808
39	1,787
40	1,576
41	1,489

Based on these results, it appears that the best tensile fatigue results were obtained with the composite formed of fibers sized with the composition of Sample 37.

5

10

EXAMPLE VIII

Four sizing compositions (samples 43-46) were prepared according to the method of Example I having the formulations described in Table 15.

TABLE 15

SAMPLE		43		44		45		46
INGREDIENTS	%	gms	%	gms	%	gms	%	gms
A1100	1.5	210	1.5	165	1.5	165	1.5	165
TA22	2.25	315	2.25	247				
NЩоН	1.4	200	1.4	154				
MB4000M ⁸	12.0	1680						
Aquathane 52-00-01	0.5	70	0.5	55	0.5	55		
Moldpro 932	2.25	315	2.25	247	3.0	330	3.0	330
PP448C			12.0	1320	11.0	1200		
PP353C							120	1320
Quantity Solids (gms)		14,000		11,000		11,000		11,000

⁸ MB4000M is a nonionic polypropylene dispersion made from medium molecular weight maleated resin converted by Michelman Inc.

These sizing compositions were used to size glass fibers and the physical strand properties were assessed as described in Example IV. The results are shown below in Table 16.

5

TABLE 16 .

SAMPLE	ST. SOLIDS (%)	FINES (%)	ST. INTEGRITY (%)
43	0.97	3.3	7.3
44	0.9	4.8	4.3
45	0.69	3.9	4.9
46	0.61	4.7	15.3

The sized strands were chopped and formed into polypropylene composites according to the method discussed above in Example I. The composite properties were assessed and are provided in Table 17.

10

TABLE 17

SAMPLE	TENSILE	FLEX.	FLEX.	NOTCHED	UNNOTCHED	GLASS
	ST	ST	MOD.	IZOD	IZOD	CONTENT
	(PSI)	(PSI)	(KSI)	(FT-LB/IN)	(FT-LB/IN)	(%)
43	12,153	18,171	691	2.167	11.833	30.2
44	12,147	18,328	691	2.114	12.465	30.4
45	12,372	17,970	649	2.342	13.044	29.8
46	11,657	18,571	700	2.272	13.325	29.6

The foregoing Examples are provided for purposes of illustrating the sizing composition of the invention, and are not intended to limit the scope of the invention as defined by the claims which follow.

WHAT IS CLAIMED IS:

20

1. A sizing composition for application to glass reinforcing fibers comprising: a maleic acid or anhydride modified polypropylene film former having a molecular weight in excess of 35,000 Daltons, an aminosilane coupling agent, a cationic lubricant, and a fatty acid nucleating agent having from 10 to 18 carbon atoms.

- 2. The composition of claim 1, wherein said modified polypropylene film former comprises polypropylene grafted with maleic anhydride.
- 3. The composition of claim 2, wherein said modified polypropylene film former has a molecular weight in excess of about 80,000.
- The composition of claim 2, wherein said cationic modified polypropylene film former is present in an amount of from about 40 to about 80 weight percent of the total dried sizing; said aminosilane is present in an amount of from about 10 to about 20 weight percent of the total dried sizing; said cationic lubricant is present in an amount of from about 0.25 to about 1.3 weight percent of the total dried sizing; and said nucleating agent is present in an amount of from about 8 to about 15 weight percent of the total dried sizing.
 - 5. The composition of claim 2, wherein said aminosilane is a mono- or di-aminated aminosilane.
 - 6. The composition of claim 5, wherein said aminosilane is gamma-aminopropyltriethoxysilane.
 - 7. The composition of claim 2, wherein said cationic lubricant is selected from the group consisting of ethoxylated amines.
 - 8. The composition of claim 7, wherein said cationic lubricant is polyethyleneinine polyamide.
- 9. The composition of claim 2, wherein said nucleating agent is selected from the group consisting of terephthalic acid, sebasic acid, stearic acid, palmitic acid, and mixtures thereof.
 - 10. The composition of claim 9, wherein said nucleating agent is terephthalic acid.
- 11. The composition of claim 2, wherein said aminosilane is a mono- or di-aminated aminosilane, said cationic lubricant is an ethoxylated amine and said nucleating agent is selected from the group consisting of terephthalic acid, sebasic acid, stearic acid, palmitic acid and mixtures thereof.

12. Glass fibers for reinforcing polymeric materials comprising, glass fibers coated with the dried residue of a sizing composition comprising: a maleic acid or anhydride modified polypropylene film former having a molecular weight in excess of about 35,000, an amino silane coupling agent, a cationic lubricant, and a fatty acid nucleating agent having 10 to 18 carbon atoms.

13. The glass fibers of claim 12, wherein said cationic polypropylene film former comprises polypropylene grafted with maleic anhydride.

5

30

- 14. The glass fibers of claim 13, wherein said modified polypropylene film former has a molecular weight in excess of about 80,000.
- 15. The glass fibers of claim 13, wherein, said cationic polypropylene film former is present in an amount of from about 40 to about 80 weight percent of the total dried sizing; said amino silane is present in an amount of from about 10 to about 20 weight percent of the total dried sizing; said cationic lubricant is present in an amount of from about 0.25 to about 1.3 weight percent of the total dried sizing; and said nucleating agent is present in an amount of from about 8 to about 15 weight percent of the total dried sizing.
 - 16. The glass fibers of claim 13, wherein said aminosilane is a mono- or di-aminated aminosilane.
 - 17. The glass fibers of claim 16, wherein said aminosilane is gamma-aminopropyltriethoxysilane.
- 20 18. The glass fibers of claim 13, wherein said cationic lubricant is selected from the group consisting of ethoxylated amines.
 - 19. The glass fibers of claim 18, wherein said cationic lubricant is polyethyleneimine polyamide.
- The glass fibers of claim 13, wherein said nucleating agent is selected from the
 group consisting of terephthalic acid, sebasic acid, stearic acid, palmitic acid and mixtures thereof.
 - 21. The glass fibers of claim 20, wherein said nucleating agent is terephthalic acid.
 - 22. The glass fibers of claim 13 wherein said aminosilane is a mono- or di-aminated aminosilane, said cationic lubricant is an ethoxylated amine and said nucleating agent is selected from the group consisting of terephthalic acid, sebasic acid, stearic acid, palmitic acid and mixtures thereof.
 - 23. A polypropylene matrix reinforced by glass fibers defined in claim 13.

24. A method for improving tensile creep and tensile fatigue in polypropylene matrix composites reinforced with glass fibers comprising treating said glass fibers with the sizing composition of claim 1 prior to their introduction into said polypropylene matrix material.

INTERNATIONAL SEARCH REPORT

Inte. ional Application No PCT/US 00/03609

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C03C25/26 C08J C08J5/08 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) C03C C08J Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) EPO-Internal C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Category ° Relevant to daim No. X US 4 426 469 A (GARAGNANI ENEA ET AL) 1-24 17 January 1984 (1984-01-17) column 2, line 13 - line 64; examples X US 4 448 917 A (BLACK DENNY E ET AL) 1-24 15 May 1984 (1984-05-15) column 1, line 26 -column 2, line 5; claims US 5 648 169 A (HOU YONGSHENG ET AL) A 1,2, 15 July 1997 (1997-07-15) 4-13 cited in the application 15-24 column 3, line 9 -column 4, line 48 A US 5 437 928 A (HOU YONGSHENG ET AL) 1-14 1 August 1995 (1995-08-01) the whole document -/--Further documents are listed in the continuation of box C. X Patent family members are listed in annex. Special categories of cited documents : "T" later document published after the international filing date or priority date and not in conflict with the application but "A" document defining the general state of the art which is not considered to be of particular relevance cited to understand the principle or theory underlying the invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) 'Y' document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-ments, such combination being obvious to a person skilled in the art. *O* document referring to an oral disclosure, use, exhibition or document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 28 June 2000 05/07/2000 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016 Reedijk, A

INTERNATIONAL SEARCH REPORT

Inter- inal Application No PCT/US 00/03609

C(Continua	tion) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
4	FR 2 735 151 A (VETROTEX FRANCE SA) 13 December 1996 (1996-12-13) the whole document	1-24
	US 5 804 313 A (SCHELL PHILIP L) 8 September 1998 (1998-09-08) column 16, line 18 - line 24; table 1	6-11, 16-22
		·
į		

INTERNATIONAL SEARCH REPORT

information on patent family members

Inte. onal Application No PCT/US 00/03609

Patent document cited in search report	t	Publication date		atent family member(s)	Publication date
· · · · · · · · · · · · · · · · · · ·		LL			<u> </u>
US 4426469	Α	17-01-1984	IT	1147319 B	19-11-1986
			DE	3170466 D	20-06-1985
			EP	0036097 A	23-09-1981
			JP	1004530 B	26-01-1989
		•	JP	1526635 C	30-10-1989
			JP	56140049 A	02-11-1981
US 4448917	A	15-05-1984	US	4489131 A	18-12-1984
US 5648169	Α	15-07-1997	US	5437928 A	01-08-1995
			CA	2174420 A	04-05-1995
			CN	1134130 A	23-10-1996
			EP	0725729 A	14-08-1996
			JP	3003705 B	31-01-2000
			JP	9504344 T	28-04-1997
			WO	9511800 A	04-05-1995
US 5437928	Α	01-08-1995	CA	2174420 A	04-05-1995
			CN	1134130 A	23-10-1996
			EP	0725729 A	14-08-1996
			JP	3003705 B	31-01-2000
			ĴΡ	9504344 T	28-04-1997
			WO.	9511800 A	04-05-1995
			ÜS	5648169 A	15-07-1997
FR 2735151	Α	13-12-1996	AU	712261 B	04-11-1999
			AU	6229496 A	09-01-1997
			BR	9606431 A	30-09-1997
			CA	2195756 A	27-12-1996
			CN	1159217 A	10-09-1997
			CZ	9700365 A	11-06-1997
			. EP	0776393 A	04-06-1997
			WO	9641912 A	27-12-1996
			HÜ	9700374 A	29-12-1997
			JP	10505141 T	19-05-1998
			NO	970199 A	16-01-1997
			PL	318535 A	23-06-1997
			SK	17497 A	09-07-1997
			TR	970349 T	22-04-1997
			US	6051316 A	18-04-2000
					10-04-2000
US 5804313	Α	08-09-1998	NONE	•	